

## Elegetolepis and its kin, the earliest monodontode chondrichthyans

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***Elegestolepis* and its kin, the earliest monodontode chondrichthyans**

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*Elegestolepis* and Its Kin, the Earliest Monodontode Chondrichthyans

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Running header—Earliest monodontode chondrichthyans

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4 20 ABSTRACT—Chondrichthyan-like scales with simple, single odontode crowns,  
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6 21 reminiscent of those of euselachians, have been reported from Silurian strata in a  
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8 22 number of previous studies. These specimens comprise the genera *Elegestolepis* (from  
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10 23 Siberia, Mongolia and Tuva) and *Kannathalepis* (from the Canadian Arctic), and have  
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12 24 been considered to exhibit contrasting patterns of ontogenetic development.  
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16 25 A study of elegestolepid microremains from the Chagat Formation of Mongolia  
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18 26 (Llandovery–lower Wenlock) and the Baital Formation of Tuva (Wenlock–Ludlow) has  
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20 27 been undertaken using SEM and micro-CT to examine scale canal system and hard  
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22 28 tissue structure. These investigations revealed scales at different stages of  
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24 29 development, whose morphogenesis is characterized by growth (elongation) of the  
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26 30 crown odontode and formation of neck canals. This ontogenetic pattern (*Elegestolepis*-  
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28 31 type morphogenesis) is also recognized in *Kannathalepis* and the Lower Devonian  
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30 32 species *Ellesmereia schultzei*, and forms the basis for the unification of these taxa into a  
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32 33 new chondrichthyan Order Elegestolepidida ordo nov. Similarities in crown  
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34 34 vascularization (branching pulp, single neck canal) shared by *Elegestolepis*,  
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36 35 *Ellesmereia* and *Deltalepis* gen. nov. (*D. magna* sp. nov. and *D. parva* sp. nov. erected  
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38 36 here in for Mongolian specimens) require the erection of the Family Elegestolepididae  
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40 37 fam. nov. that is distinguished from the mono-generic Kannathalepididae (non-  
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42 38 branching pulp, multiple neck canals).  
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50 39 Elegestolepid scales exhibit characteristics (neck canal formation and lack of enamel  
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52 40 and basal bone osteons) consistent with those of the chondrichthyan dermal skeleton.  
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55 41 This establishes Elegestolepidida as the stratigraphically oldest chondrichthyan taxon to  
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3 42 develop monodontode scales, which, in contrast to the ‘placoid’ scales of euselachians,  
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6 43 are growing structures.  
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## INTRODUCTION

The type species of the genus *Elegestolepis* (*E. grossi*) was described by Karatajūtė-Talimaa (1973) from isolated scales from upper Ludlow–Pridoli strata (Elegest and Kadvoj outcrops, Tuva, Russian Federation) of the Tuva-Mongol terrane and, at the time of publication, was the earliest known taxon referred to the Chondrichthyes. Subsequent studies on microvertebrate fossils from the lower Paleozoic have led to the identification of stratigraphically older species attributed to *Elegestolepis*. These are represented by middle Llandovery–lower Wenlock *E. sp.* specimens from the Tuva-Mongol (Chargat Formation, north western Mongolia; Elegest Formation, Tuva, Russia (Karatajūtė-Talimaa et al., 1990; Sennikov et al., 2015)) and Altai terranes (Gornaya Shoriya, Altai Republic, Russia (Sennikov et al., 2015)) and the middle–upper Llandovery *E. conica* from the adjacent Siberian craton (Angara-Ilim, Niuya-Bresovo and Tchuna-Biriussa sections, Siberian District, Russia (Karatajūtė-Talimaa and Predtechenskyj, 1995)). The paleogeographical and stratigraphical range of *Elegestolepis*-like taxa was further expanded with the description (Vieth, 1980) of the Laurussian chondrichthyan scale species *Ellesmereia schultzei* (from the Lochkovian of Ellesmere Island, Nunavut Territory, Canada).

According to the categories of scale morphogenesis established by Karatajūtė-Talimaa (1992) for Paleozoic chondrichthyans, *Elegestolepis* and *Ellesmereia* belong to the *Elegestolepis* developmental type as the scales have a monodontode, non-growing crown enclosing a pulp canal that opens at the crown neck via a single foramen. Influenced by the lepidomorial theory put forward by Stensiö and Ørvig (Stensiö and

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3 84 Ørvig, 1951–1957; Stensiö, 1961), Karatajūtė-Talimaa (Karatajūtė-Talimaa, 1992;  
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6 85 Karatajūtė-Talimaa, 1998) proposed that elegestolepid scale crowns represent the  
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8 86 simplest monodontode dermatoskeletal elements, exhibiting many of the characteristics  
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10 87 of what were assumed to be the most elementary skeletal units of the integument  
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12 88 (lepidomoria). Thus, the development of odontodes in elegestolepids was differentiated  
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15 89 from other chondrichthyans with ‘placoid’ (monodontode) scales, where the crowns  
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18 90 were considered to form through the coalescence of lepidomoria. Karatajūtė-Talimaa  
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20 91 (1992, 1998) attributed this complex morphogenetic pattern to the *Polymerolepis* and  
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22 92 *Heterodontus* (euselachian; Fig. 1C) scale types. A hypothesis of odontode evolution in  
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24 93 stem chondrichthyans was founded upon these assumptions, placing lepidomorium-like  
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27 94 elements as the phylogenetic precursors of all chondrichthyan scales (Karatajūtė-  
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29 95 Talimaa, 1992).

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32 96 In the years following the conceptualization of the lepidomorial theory, increasing  
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34 97 evidence from studies on the development of the integumentary skeleton of Recent  
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37 98 neoselachians (Reif, 1980b; Miyake et al., 1999; Johanson et al., 2008) has discredited  
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39 99 the concrescence model of odontode morphogenesis, and this is now refuted by most  
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42 100 authors (Smith and Coates, 1998; Donoghue, 2002 and references therein). The latter  
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44 101 view is strengthened with the inclusion of ‘acanthodians’ bearing polyodontode scales  
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47 102 with elaborately branching odontode pulps (e.g. in *Poracanthodes* Gross, 1956;  
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49 103 Valiukevičius, 1992) within the chondrichthyan phylogenetic history (Zhu et al., 2013;  
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51 104 Brazeau and Friedman 2015; Giles et al., 2015). This contradicts the evolutionary  
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53 105 scenario predicted by the concrescence model, which claims origination of  
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56 106 neoselachian placoid scales and their complex canal system via fusion of simple  
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lepidomorial elements enclosing a single vascular loop (Stensiö and Ørvig, 1951–1957; Stensiö, 1961). Advances in developmental biology have revealed an apparently conserved gene regulatory network that maintains a variety of odontode morphogenetic pathways (Fraser et al. 2010). This further corroborates the notion that all structures resolvable into odontode units are, in a broad sense, homologous. In this context, a re-examination of *Elegestolepis* and *Elegestolepis*-like Silurian scale taxa (e.g. *Ellesmereia*, *Kannathalepis*) will enable a clearer understanding of the early evolution of single odontode integumentary skeletal elements in the Chondrichthyes. To meet this end, the present study investigates the histology, canal system and inferred development of *Elegestolepis grossi* scales and that of previously undescribed scales from the Lower Silurian of Mongolia referred to *Elegestolepis* (Karatajūtė-Talimaa et al., 1990). These new data permit a new systematic framework for *Elegestolepis*-like taxa and allow for the further evaluation of their likely chondrichthyan affinities.

## MATERIALS AND METHODS

The isolated scales were extracted through dissolution of carbonate rock samples with dilute acetic acid. The specimens come from the Chargat Formation of north western Mongolia (Chargat outcrop, sample P-16/3 [2]) and the Baital Formation (Elegest River outcrop, samples from beds 236, 291, 293 and 295 [1]) of central Tuva, Russian Federation.



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3 128 Scale morphology was documented using the Zeiss EVO LS and the JEOL JSM-  
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5 129 6060 scanning electron microscopes at the School of Dentistry of the University of  
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8 130 Birmingham, UK. Prior to imaging, the specimens were sputter-coated with a 25 nm-  
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10 131 thick layer of gold/palladium alloy.  
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13 132 Hard tissue microstructure and internal architecture of thin-sectioned specimens was  
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15 133 investigated by Nomarski differential interference contrast microscopy (using a 'Zeiss  
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17 134 Axioskop Pol' polarization microscope) and scanning electron microscopy (with a JEOL  
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19 135 JSM-6060 SEM at the School of Dentistry, University of Birmingham, UK).  
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23 136 Scale examination with X-ray radiation was conducted using the SkyScan 1172  
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25 137 microtomography scanner at the School of Dentistry of the University of Birmingham,  
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27 138 UK. The acquired microradiographs (tomographic projections) were taken at 0.3°  
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29 139 intervals over a 180° rotation cycle at exposure times of 400 ms, using a 0.5 mm thick  
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31 140 X-ray attenuating Al filter. These image data were processed with the SkyScan NRecon  
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33 141 reconstruction software in order to generate sets of microtomograms that were  
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35 142 converted into volume renderings in Amira 5.4 3D analysis software.  
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39 143 Figured material is deposited in the Lapworth Museum of Geology, University of  
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41 144 Birmingham, UK (BU prefix).  
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49 **Definitions of Terms**  
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52 147 Traditionally (Sykes, 1974; Duffin and Ward, 1993; Thies, 1995) the two main  
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54 148 components (crown and base) of chondrichthyan scales have been identified on the  
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56 149 basis of morphological and/or topological criteria without consideration of their  
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developmental origin. This approach can lead to ambiguity when attempting to establish the extent of these structures and, more importantly, can result in homologizing scale parts with different tissue composition across taxa. To address the above issues, Andreev et al. (2015) provided revised definitions of terms used in literature to describe chondrichthyan scales, and these are followed in the present study.

## SYSTEMATIC PALEONTOLOGY

Class CHONDRICHTHYES Huxley, 1880

Order ELEGESTOLEPIDIDA ordo nov.

**Included Families**—Kannathalepididae Märss and Gagnier 2001 and Elegestolepididae fam. nov.

**Diagnosis**—Chondrichthyan fish with monodontode scale crowns composed of a growing odontode that encloses neck-canal branches of the pulp cavity (Fig. 1B).

**Remarks**—The recent literature on putative basal chondrichthyan taxa (e.g. mongolepids, elegestolepids, kathermacanthids and polymerolepidiforms) from the lower Paleozoic expresses uncertainty regarding their systematic position relative to the major clades (Subclasses) of the Chondrichthyes (Karatajute-Talimaa and Novitskaya, 1997; Sansom et al., 2000; Märss et al., 2006; Hanke and Wilson, 2010; Hanke et al., 2013). This reflects an inadequate understanding of the phylogenetic significance of scale-

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3 171 derived characters, which have been employed to diagnose these taxa given the  
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5 172 general absence of chondrichthyan endoskeletal and dental remains in the lower  
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8 173 Paleozoic.  
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11 174 The odontode growth that typifies the ontogenesis of *Elegestolepis*-like scales is  
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13 175 not seen within traditionally recognised chondrichthyan clades (*sensu* Grogan et al.,  
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15 176 2012), yet the *Elegestolepidida* consistently falls inside stem-group Chondrichthyes  
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17 177 when its affinities are tested via phylogenetic inference (Andreev et al., unpublished  
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19 178 data; Fig. 2). The erection of a new Order draws together species that possess scales  
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21 179 with growing single-odontode crowns whose morphogenesis can be differentiated from  
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23 180 that of elasmobranch ‘placoid’ scales (the *Heterodontus* morphogenetic type of  
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25 181 (Karatajūtė-Talimaa, 1992, 1998; Fig. 1C). The formal recognition of the *Elegestolepis*-  
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27 182 type of scale development represents a change in concept from what was originally  
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29 183 identified as a purely morphogenic category (Karatajūtė-Talimaa, 1992, 1998).  
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39 184  
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42 185 Family KANNATHALEPIDIDAE Märss and Gagnier, 2001  
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45 186 **Included genera**—*Kannathalepis* Märss and Gagnier, 2001  
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48 187 **Revised diagnosis**—*Elegestolepids* possessing dermal scales with vertically  
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50 188 undivided pulp cavities from which multiple (up to five) horizontal neck canals emerge  
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52 189 basally.  
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54 190 **Remarks**—The mono-generic Family Kannathalepididae was introduced by Märss  
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56 191 and Gagnier (2001) to distinguish *Kannathalepis*, identified to exhibit a specialised type  
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3 192 of scale morphogenesis, from other Silurian chondrichthyan scale taxa (mongolepid and  
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6 193 elegestolepid). It was reported that the squamation of *Kannathalepis* consists of single-  
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8 194 odontode scales along with more complex aggregates of fused 'placoid' scales that  
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10 195 were thought to provide evidence for two separate modes of scale development within  
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12 196 the genus (Märss and Gagnier, 2001: fig. 4f). The current study regards these  
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14 197 compound scales of *Kannathalepis* as aberrant, formed by anomalous patterning that is  
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16 198 thought to result from suppression of inter-scale domains in accordance with the  
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18 199 inhibitory field model outlined by Reif (1980a, 1982). Localised suturing of scales has  
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20 200 similarly been documented in stem (*Hybodus delabechei* (Reif, 1978: fig. 8 d, e) and  
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22 201 *Lissodus sardiniensi* (Fischer et al., 2010: fig. 7l)) and crown (*Echinorhinus brucus* (Reif,  
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24 202 1985:pl. 15) and *Asterodermus platypterus* (Thies and Leidner, 2011:pl. 71))  
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26 203 euselachians with monodontode trunk scale cover that is known to be prevalent within  
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28 204 the Order (Reif, 1985; Thies and Leidner, 2011; Dick, 1978; Dick and Maisey, 1980;  
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30 205 Maisey, 1989; Wang et al., 2009).

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37 206 Complexes of randomly sutured monodontode scales consequently cannot be  
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39 207 considered equivalent to polyodontode scales (e.g. those of Mongolepidida (Karatajūtė-  
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41 208 Talimaa, 1998)), since the odontodes of the latter are patterned as a unit in a particular  
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43 209 manner and are given support by a common base/pedicle tissue. The scale  
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45 210 development in *Kannathalepis* can thus be identified as that of 'placoid' scales with a  
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47 211 growing odontode and base, corresponding to the *Elegestolepis* morphogenetic type  
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49 212 (Fig. 1B) of Karatajūtė-Talimaa (1992). On that basis, Kannathalepididae is placed  
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51 213 inside the new Order Elegestolepidida, and its validity is maintained by acknowledging  
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the canal system characteristics (vertically undivided pulp cavity and multiple neck  
canals) diagnostic for the Family, recognized in the original description of the taxon.

Kannathalepididae was expanded subsequent to its erection to include the  
Wenlockian genus *Frigorilepis*, which was described from articulated body fossils  
(Märss et al., 2002, 2006). Nevertheless, crown morphogenesis in *Frigorilepis* has not  
been demonstrated to proceed in discrete growth phases as in elegestolepid taxa,  
which are further distinguished by the presence of scale-neck canal openings. The  
absence of characters diagnostic for Elegestolepidida results in treating *Frigorilepis* as  
Family and Order *incertae sedis* for the time being.

Family ELEGESTOLEPIDIDAE fam. nov.

**Included genera**—The type genus *Elegestolepis* Karatajūtė-Talimaa 1973,  
*Ellesmereia* Vieth, 1980 and *Deltalepis* gen. nov.

**Diagnosis**—Elegestolepids with scales that develop a vertically branched pulp cavity  
that gives off a single horizontal neck canal and dentine canals that originate at the  
lower neck/pedicle surface independently of the pulp (Fig. 9).

Genus *ELEGESTOLEPIS* Karatajūtė-Talimaa, 1973

**Included species**—The type species *E. grossi* Karatajūtė-Talimaa, 1973 and *E.*  
*conica* Novitskaya and Karatajūtė-Talimaa, 1986.

**Revised diagnosis**—*Elegestolepidids* possessing up to three unornamented scale crown lobes (Fig. 3A; Fig. 4A, B, D, E) incised by deep, linear grooves.

*ELEGESTOLEPIS GROSSI* Karatajūtė-Talimaa, 1973  
(Figs. 1B, 3A, 4, 5, 9A–C)

*Elegestolepis grossi* Karatajūtė-Talimaa, 1973:figs. 1–5, pl. 3 (original description).

*Elegestolepis grossi* Karatajūtė-Talimaa, 1998:31, fig. 10.

**Locality and horizon**—Studied material comes from beds 236, 291, 293 and 295 of the Baital Formation (Wenlock–Ludlow (Vladimirskaya, 1978; Sennikov et al., 2015)) at the type locality on the Elegest River, central Tuva, Russia (Karatajūtė-Talimaa, 1973). *E. grossi* has also been reported from the Pridoli of Tuva, Russia (Khondergei Formation (Sennikov et al., 2015)) as well as from strata of the lower Wenlock Upper Tarkhata Subformation (Gorny Altai, Russia (Sennikov et al., 2015)).

**Holotype**—An ontogenetically mature scale (T-003) from the Baital Formation of Tuva, Russian Federation (Karatajūtė-Talimaa, 1973).

**Referred material**—Over 200 isolated scales that were examined for this study are deposited in the Lapworth Museum of Geology, University of Birmingham, UK.

**Revised diagnosis**—*Elegestolepis* species possessing small (up to c. 1 mm long) scales that have deltoid to lanceolate, trilobate crowns and develop moderately to strongly constricted necks and bulbous bases during their ontogenesis. Scale odontode composed of dentine tissue with multipolar odontocyte lacunae from which emerge

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3 255 canaliculi with dendroid branching. Cellular basal bone with layered mineralised-fibre  
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6 256 organization.  
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9 257       **Remarks**—Certain differences were noted between the scale histology of *E.*  
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11 258 *grossi* scales and the original descriptions of Karatajūtė-Talimaa (1973). Some of these  
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13 259 concern the nature of the most superficial portion of the scale crown and neck,  
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16 260 understood by Karatajūtė-Talimaa (1973) to consist of a type of hypermineralized  
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18 261 dentine tissue, durodentine (one of the less commonly used synonyms of enameloid  
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20 262 (Ørvig, 1967; Smith and Miles, 1971; Sire et al., 2009). This ‘enameloid’ layer is found  
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22 263 not to be a persistent feature of *E. grossi* scales, and even when present it appears  
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24 264 discontinuous across most of the upper crown surface (Fig. 5A–E), contrary to previous  
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26 265 depictions (Karatajūtė-Talimaa, 1973: fig. 2a, b and Sire et al., 2009: fig. 10b). The layer  
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28 266 is instead most prominent around the scale neck (Fig. 5A, C–E) and can extend all the  
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30 267 way down to the level of the basal bone (Fig. 5C). This distribution is contrary to that of  
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32 268 single crystalline enameloid in neoselachian scales, where it is confined mainly to the  
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34 269 upper crown region (Johns et al., 1997). Furthermore, the architecture of the superficial  
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36 270 crown region cannot be recognised in any of the known enameloid structural types  
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38 271 (Johns et al., 1997; Sansom et al., 2005; Gillis and Donoghue, 2007; Guinot and  
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40 272 Cappetta, 2011; Andreev and Cuny, 2012), but instead resembles that of the crown  
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42 273 dentine and is regarded as such. The more porous appearance of the surface dentine is  
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44 274 likely to be diagenetically induced and/or due to alteration of the original tissue  
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46 275 microstructure by preparation of the specimens with unbuffered acetic acid (even in low  
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48 276 concentration, the latter has been shown to damage the phosphatic tissues of conodont  
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50 277 elements (Jeppsson et al., 1985; Jeppsson and Anehus, 1995).  
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This study also demonstrates the presence of not previously identified faint depositional lines (Fig. 5G) in the basal bone of *E. grossi* scales, although growth of the bone tissue has been inferred from specimens in different stages of development (Karatajūtė-Talimaa 1973, 1998). The lamellae, demarcated by the depositional lines produced by change in orientation of the matrix's crystalline fibres, have convex down profiles common for the scale bases of lower vertebrates (Ørvig, 1966; Zangerl, 1968; Denison, 1979; Burrow and Turner, 1998, 1999; Qu et al., 2013).

Genus *ELLESMEREIA* Vieth, 1980

**Included species**—*Ellesmereia schultzei* Vieth, 1980

**Remarks**—*Ellesmereia* (Fig. 3B) was assigned to the Elasmobranchii by Vieth (1980) despite being recognized to possess an *Elegestolepis*-type of scale morphogenesis (Reif, 1978; Karatajūtė-Talimaa, 1992) that is atypical for an elasmobranch, and consequently it is placed here within the Elegestolepidida. Mature *Ellesmereia* scales also possess a canal system architecture (Vieth, 1980: fig. 26) closely resembling the vascularization of *Elegestolepis* and *Deltalepis* gen. nov., and for these reasons the three taxa are united at a familial level.

Genus *DELTALEPIS* gen. nov.

**Included species**—*Deltalepis magna* gen. et sp. nov. (type species) and *Deltalepis parva* gen. et sp. nov.



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**Derivation of name**—From ‘delta’ (alluding to the resemblance of the scale crown to the Greek letter Δ) and ‘lepis’, scale in Greek.

**Diagnosis**—Elegestolepidids whose scales possess lobed crowns ornamented by tuberculate ridges. Crown lobes and furrows extend down the anterior face of the scale neck (Figs. 7, 8).

**Remarks**—The material referred here to *Deltalepis* gen. nov. has not previously been formally described or figured, although was considered to belong to the genus *Elegestolepis* by Karatajūtė-Talimaa et al. (1990) and Karatajute-Talimaa and Novitskaya (1997) in their work on the mongolepid taxa from the Chargat Formation. *Deltalepis* gen. nov. scales possess crown morphology, ornamentation and pulp cavity branching pattern that differentiate them from *Elegestolepis* and *Ellesmereia*, and therefore require the erection of a new taxon. This distinction and the erection of two *Deltalepis* species is based on the documented intra- and inter-generic variation of trunk-scale morphology (e.g. crown shape, number of crown ridges/lobes and ornamentation) in Recent neoselachian Families (Reif, 1985; Compagno, 1988; Voigt and Weber, 2011). Comparable differences in ornament have also been used to distinguish taxa at genus level among thelodonts (e.g. *Erepsilepis* (Märss et al., 2006)) and mongolepid chondrichthyans (*Shiqianolepis* and *Rongolepis* (Sansom et al., 2000)). Ridged lobes are also a feature in the putative chondrichthyan taxon *Areyongalepis oervigi* (Young, 1997, 2000) known solely from micro-remains from the Darriwilian Stokes Siltstone (Amadeus Basin, Northern Territory, Australia). The crown necks and bases of elegestolepid scales, however, are not developed in *Areyongalepis* elements,

and the latter do not demonstrate identifiable vertebrate mineralised tissues (Young, 1997), making their systematic position uncertain for the time being.

*DELTALEPIS MAGNA* sp. nov.

(Figs. 3C, 6, 8A–B, 9D–F)

**Derivation of name**—From the feminine form of the Latin word for large, referring to the scale size of the species relative to that of *D. parva* gen. et sp. nov.

**Locality and horizon**—The type and only known locality for *D. magna* is 80 km north of Lake Khar-Us, north-western Mongolia (Karatajūtė-Talimaa et al., 1990). All specimens come from sample P-16/3 collected from the upper Llandovery–lower Wenlock (Salhit regional Stage) horizons of the Chargat Formation (Ørvig, 1977; Karatajūtė-Talimaa et al., 1990).

**Holotype**—An isolated, presumably trunk, scale BU5269 (Figs. 3C, 6A–C).

**Referred material**—Six isolated scales (BU5269–BU5274).

**Diagnosis**—*Deltalepis* species possessing scales with deltoid to elliptic crowns divided into three to five discrete lobes by posteriorly widening grooves. Parallel tuberculate ridges developed on the undersurface of the crown. The rami of the pulp cavity formed inside the scale crown connect directly to the main pulp canal.

**Description**

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**Morphology**—Scales possess monodontode crowns with ovate to acuminate outlines (Fig. 6) that are 500–700  $\mu\text{m}$  long and 400–700  $\mu\text{m}$  wide. The crown surface displays a complex topography that is produced by three to five lobes separated by deeply recessed inter-lobe regions (Fig. 6A–C, E, G, H). The lobes are lanceolate-shaped and can exhibit slight divergence towards the posterior of the scale. Their surface is ornamented by sub-parallel tuberculate ridges (up to 8 per lobe) that are absent from the smooth-faced inter-lobe segments of the crown. Longitudinally directed ridges are similarly developed on the undersurface of the crown (Fig. 6F, I, J), and these demonstrate regular spacing across its width.

The crown transitions into an unornamented narrow neck (down to a third of the maximal crown width) that is located at the anterior of the scale, overhung on all sides by the crown. The lower portion of the neck is either gently curved outwards or flares out to form an ellipse-shaped pedicle. In specimens with a developed pedicle support (Fig. 6E–G, I, J) the posterior face of the neck is pierced by a single centrally positioned foramen (Fig. 6F) with a diameter of c. 30–40  $\mu\text{m}$ . The lower pedicle surface of some specimens is deeply indented (Fig. 6I) and penetrated by the scale’s canal system, whereas in others it is nearly flat (Fig. 6J), exhibiting only a greatly constricted opening of the pulp.

**Histology**—The scale odontodes are composed solely from a highly vascular tubular dentine (Fig. 8A, B). The canaliculi of the dentine have a coiled appearance and display a tangled organization as well as extensive ramification along their length (up to c. 20  $\mu\text{m}$ ). In the upper portion of the crown, the canalicular network emerges from a complex of horizontally and vertically branched, interconnected, small-calibre dentine

364 canals (diameter of c. 5–25  $\mu\text{m}$ ; Fig. 9D). The latter are most prominent inside the  
365 crown lobes where they associate with and connect to branches (c. 30–60  $\mu\text{m}$  in  
366 diameter) of the pulp canal. For most of their length the pulp branches extend parallel  
367 the crown surface, before curving basally to merge (Fig. 9F) into a single pulp canal (c.  
368 60–90  $\mu\text{m}$  wide) inside the scale neck. From the posterior of the pulp issues an  
369 unbranched horizontal canal (c. 70  $\mu\text{m}$  long; Fig. 9F) that opens on the scale neck  
370 surface. Separate from the pulp cavity system, the posterior half of the scales houses  
371 numerous closely spaced (up to c. 10  $\mu\text{m}$  apart) dentine canals (10–20  $\mu\text{m}$  in diameter)  
372 whose paths parallel that of the lower crown surface (Fig. 9E). The lower ends of these  
373 canals ramify inside the scale neck before either exiting the scale basally (Fig. 9E) or  
374 ending blindly inside it.

375 The tissue (c. 40  $\mu\text{m}$  thick) closing off the lower pedicle opening displays an  
376 optically discernable boundary with the overlying dentine (Fig. 8A), but it could not be  
377 ascertained whether it constitutes a distinct tissue type.

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379 *DELTALEPIS PARVA* sp. nov.

380 (Figs. 3D, 7, 8C–D, 9G–J)

381 **Derivation of name**—From the feminine form of the Latin word for small,  
382 referring to the scale size of the species relative to that of *D. magna* gen. et sp. nov.

383 **Locality and horizon**—The type and only known locality situated 80 km north of  
384 Lake Khar-Us, north-western Mongolia (Karatajūtė-Talimaa et al., 1990). All specimens  
385 come from the upper Llandovery–lower Wenlock (Salhit regional Stage) horizons

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3 386 (sample P-16/3) of the Chargat Formation (Karatajūtė-Talimaa et al., 1990; Žigaitė et  
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5 387 al., 2011).

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8 388 **Holotype**—An isolated, presumed trunk, scale BU5275 (Figs. 3D, 7A, B).

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11 389 **Referred material**—Six isolated scales (BU5275, BU5277, BU5278–BU5280,  
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13 390 BU5282).

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17 391 **Diagnosis**—*Deltalepis* species with ovoid scale crowns compartmentalized into  
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19 392 seven to ten lobes. The lateral crown branches of the pulp cavity do not connect directly  
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21 393 to the main pulp canal.

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28 395 **Description**

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31 396 **Morphology**—The scale crowns are single odontode structures with ovoid  
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33 397 outlines (Fig. 7) that are 200–500 µm long and 200–400 µm wide. Upper crown surface  
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35 398 is divided into seven to ten antero-posteriorly aligned lobes (40–60 µm wide; Fig. 7A–F)  
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37 399 separated by much narrower, deeply incised grooves that expand towards the posterior  
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39 400 (up to c. 20 µm wide). Tubercles organized into parallel rows ornament the upper  
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41 401 surface of the crown lobes (up to three rows per lobe), whereas all other scale surfaces  
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43 402 are smooth.

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48 403 The anterior of the crown is constricted into a vertically orientated neck that  
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50 404 reaches a third to three-quarters of the maximal crown width, and which in some  
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52 405 specimens expands basally to form a pedicle support (Fig. 7C–F, H, I). The posterior  
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54 406 lower-neck/pedicle face of these scales is pierced by a single foramen (Fig. 7D, H, I)

with a diameter of 20–35  $\mu\text{m}$ . A canal opening is also present on the lower pedicle surface (Fig. 7H), while a row of elliptical foramina of laterally decreasing diameter (from 70  $\mu\text{m}$  to 40  $\mu\text{m}$  in Fig. 7G) mark the lower face of scales lacking a pedicle attachment.

**Histology**—Tubular dentine tissue (Fig. 8C, D) is the only component of the scale crown. The dentine canaliculi are less than 2  $\mu\text{m}$  in diameter and up to c. 20  $\mu\text{m}$  long, with arborescent branching (Fig. 8D) that gives the tubular system a tangled appearance. Inside the lobed regions of the crown, the tubules connect to a network of vertically (c. 5–10  $\mu\text{m}$  wide and 25–40  $\mu\text{m}$  long) and horizontally (c. 5  $\mu\text{m}$  wide) oriented dentine canals (Fig. 8C; Fig. 9J) that are confluent with branches of the pulp cavity. These pulp branches (from c. 20  $\mu\text{m}$  to c. 45  $\mu\text{m}$  in diameter; Fig. 9G–J) occupy the crown lobes (one canal per lobe) before curving basally to merge with one another inside the scale neck. The three medial branches emerge from the main pulp canal—confined to the scale neck/pedicle—whereas the more lateral ones are only indirectly connected to it through the medial rami (Fig. 9I). Near its lower end, the main pulp canal gives off a short neck canal (Fig. 9I, J) that opens at the scale surface.

Posterior of the pulp-cavity canal system the scale houses a number (c. 15) of mutually parallel, ascending dentine canals (Fig. 9G) with diameters between c. 10  $\mu\text{m}$  and 15  $\mu\text{m}$ . These canals follow the posterior scale profile without establishing connections at any point with the pulp cavity and terminate basally at the lower pedicle surface.

## DISCUSSION

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430     **Chondrichthyan Characteristics of Elegestolepid Scales**

431             The odontogenic component of the vertebrate skeleton develops primarily as  
432 discrete elements (odontodes), each of which being the product of a single epithelia-  
433 mesenchymal cell condensation (Ørvig, 1977; Reif, 1982; Fraser et al., 2010).  
434 Odontodes are one of the main structural units of scales and in certain groups (e.g. in  
435 neoselachian chondrichthyans (Fig. 1C; Sire and Huysseune, 2003; Eames et al., 2007;  
436 Sire et al., 2009) can form the entire squamation in the absence of osteogenic  
437 contribution to the integumentary skeleton. In lower Paleozoic vertebrates, dermal  
438 odontodes are usually patterned in clusters (polyodontodia in Ørvig, 1977) that form  
439 compound scale crowns; these have been documented in pteraspidomorphs (Gross,  
440 1961; Denison, 1967; Sansom et al., 2009; Keating et al., 2015), anaspids (Märss,  
441 1968; Blom et al., 2002; Märss, 2002; Keating and Donoghue, 2016), galeaspids (Wang  
442 et al., 2005), osteostracans (Stensiö, 1932; Märss et al., 2014) and jawed  
443 gnathostomes (Schultze, 1968, 1977; Gross, 1969; Denison, 1979; Karatajūtė-Talimaa,  
444 1995; Sansom et al., 1996; 2012; Burrow and Turner, 1998, 1999; Giles et al., 2013).  
445 The Thelodonti (Märss et al., 2007), Elegestolepidida (Karatajūtė-Talimaa, 1973, 1998  
446 and this study) and some euchondrichthyans (*sensu* Janvier and Pradel, 2015)—e.g.  
447 iniopterygians (Zangerl R, Case, 1973; Grogan and Lund, 2009), petalodonts (Malzahn,  
448 1968), symmoriiforms (Lund, 1985, 1986; Coates and Sequeira, 2001), living  
449 holocephalians (Patterson, 1965) and euselachians (Thies and Leidner, 2011)—are the  
450 exception, as their scale crowns form only from a single-odontode element.

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3 451 The integumentary skeleton of thelodonts demonstrates perhaps the most  
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6 452 phylogenetically primitive mode of monodontode scale morphogenesis (Figs. 1A, 2; Sire  
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8 453 et al., 2009; Smith and Hall, 1990, 1993). In contrast to polyodontode scale  
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10 454 development, where each of the component odontodes mineralizes in a single step, the  
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12 455 scales of thelodonts go through several ontogenetic phases that result in gradual  
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14 456 elongation of the crown in basal direction (Gross, 1967; Karatajūtė-Talimaa, 1978).  
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17 457 Thelodonts can also possess basal bone tissue (Fig. 1A), the deposition of which  
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19 458 commences only after cessation of odontode growth (Karatajūtė-Talimaa, 1978; Märss  
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21 459 et al., 2007). It is argued here that a thelodont-like pattern of scale development evolved  
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23 460 convergently in the early chondrichthyans (Fig. 2), with the appearance of  
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25 461 Elegestolepida in the middle Llandovery. Nevertheless, during ontogenesis  
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27 462 elegestolepid scales develop a more derived canal system architecture that features  
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29 463 neck canal opening(s) of the odontode pulp (documented outside the Euchondrichthyes  
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31 464 in 'acanthodians' (Denison, 1979) and stem osteichthyans (Gross, 1953, 1968; Qu et  
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33 465 al., 2013) but absent from the dermal skeleton of the Thelodonti (Fig. 2; Gross, 1967;  
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35 466 Karatajūtė-Talimaa, 1978; Märss et al., 2007). The depth of insertion of the scale into  
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37 467 the integument has been suggested to influence the formation of neck canals (Hanke  
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39 468 and Wilson, 2010) and this interpretation is supported by the position of scale necks  
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41 469 inside the upper vascular layer (stratum spongiosum) of the dermis in Recent  
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43 470 neoselachians (Reif, 1980b; Miyake et al., 1999). Similar topological relationship  
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45 471 between scales and surrounding integumentary tissues is attributed here to the  
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47 472 elegestolepids, whereas the dermal odontode papillae of thelodonts have been  
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3 473 interpreted to form superficially at the epithelium-mesenchyme boundary and therefore  
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6 474 not to intersect the vascular system (Karatajūtė-Talimaa, 1978; Märss et al., 2007).  
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9 475 Outside the Chondrichthyes, other derived gnathostomes regarded to possess  
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11 476 monodontode body scales belong to the basal ‘placoderm’ Orders Stensioellida and  
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13 477 Antiarcha (Fig. 2; also refer to Johanson, 2002; Giles et al., 2015; Brazeau and  
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16 478 Friedman, 2015 and citations therein for recent vertebrate phylogenies) whose scale  
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18 479 structure is still insufficiently investigated. The available data on the squamation of  
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20 480 these taxa (e.g. *Stensioella* (Gross, 1962) and *Parayunnanolepis* (Upeniec, 2011; Zhu  
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22 481 et al. 2012)) provides evidence for non-growing odontodes, implying this to be a  
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25 482 plesiomorphic feature of the single-odontode scales of jawed gnathostomes.  
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28 483 Histological descriptions of scale hard-tissues are presently not available for the above  
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30 484 taxa, but known examples of ‘placoderm’ scale structure often demonstrate formation of  
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32 485 an osteon-rich vascular layer inside the upper portion of the basal bone (Burrow and  
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35 486 Turner 1998, 1999; Giles et al., 2013; Rücklin and Donoghue 2015). Osteon mediated  
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37 487 bone remodeling and resorption is widespread in the dermal skeleton of ‘placoderms’  
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39 488 (Donoghue et al. 2006; Downs and Donoghue, 2009; Giles et al., 2013) and basal  
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42 489 osteichthyans (Zhu et al., 2006), but, critically, is absent from the elegestolepid skeleton  
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44 490 and in conventional chondrichthyans. Other characteristics placing elegestolepids with  
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46 491 the Chondrichthyes among derived gnathostomes are the pattern of scale histogenesis  
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48 492 and their hard tissue composition, both of which match those of polyodontode  
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51 493 chondrichthyan scales by being two-component skeletal elements formed out of  
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54 494 lamellar basal bone and crown dentine (Karatajūtė-Talimaa, 1992).  
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## Elegestolepidida in the Context of Paleozoic Chondrichthyans

Although rare, elegestolepids are a significant component of pre-Devonian chondrichthyan faunas with five currently identified species grouped into two Families (Fig. 10), being second only in diversity to the Order Mongolepidida (Karatajūtė-Talimaa et al., 1990; Karatajute-Talimaa and Novitskaya, 1992, 1997; Sansom et al., 2000, 2001). Whilst the mongolepids (Sansom et al., 2001) and several other putative chondrichthyan lineages (represented by *Areyongalepis* (Young, 1997), *Tantalepis* (Sansom et al., 2012), *Tezakia* (Sansom et al., 1996; Andreev et al., 2015) and *Canyonlepis* (Sansom et al., 2001; Andreev et al., 2015)) have their origination in the Ordovician, no remains attributable to Elegestolepidida have yet to be reported from this period. These Ordovician taxa possess compound (polyodontode) scale crowns and lack neck canal openings; the latter are now understood not to develop in all basal chondrichthyans (Märss et al., 2007; Hanke and Wilson, 2010).

Neck pulp-canal openings stratigraphically first appear in the oldest elegestolepid species (*E. conica* Novitskaya and Karatajūtė-Talimaa, 1986; Karatajūtė-Talimaa and Predtechenskyj, 1995), in the Middle Llandovery, and can be recognized as a persistent feature of the canal system of mature elegestolepid scales (Fig. 10; Karatajūtė-Talimaa, 1973; Vieth, 1980; Märss and Gagnier, 2001). This condition is similarly developed in Silurian polyodontode chondrichthyan species (e.g. *Tuvalepis*, Žigaitė and Karatajūtė-Talimaa, 2008) and the monogolepids *Mongolepis*, *Teslepis* (Karatajūtė-Talimaa, 1998), *Shiqianolepis* and *Rongolepis* (Sansom et al., 2000). In monogolepids pulps exit the lower part of crown either by giving off short rami (termed 'horizontal canals' by Karatajūtė-Talimaa (1995) and considered equivalent to the neck canals of

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3 519 elegestolepid scales) or opening directly to the crown surface (in *Shiqianolepis* and  
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6 520 *Rongolepis* (Sansom et al., 2000)).  
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9 521 Elegestolepidida and Mongolepidida might represent two distinct lineages of  
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11 522 early chondrichthyans that provide an insight into the variability of scale characteristics  
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13 523 within what appear to be monophyletic groups. Inside each of these clades the features  
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16 524 shared by its member genera are those relating to the pattern of crown morphogenesis,  
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18 525 whereas aspects of their scale vascularization and hard tissue structure can exhibit  
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20 526 differences. Moreover, characters with a limited distribution in one of the Orders can  
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23 527 have a constant presence in the other, as is the case with the neck canal openings of  
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25 528 the elegestolepids. The identification of elegestolepid taxa is thus regarded to require  
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28 529 the unique character combination of a growing monodontode scale crown (Order-grade  
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30 530 character) and neck canal openings (plesiomorphy of crown-group gnathostomes).  
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33 531 Under the diagnosis formulated here, the Wenlockian species *Frigorilepis*  
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35 532 *caldwelli*, placed inside Kannathalepididae by Märss et al. (2002, 2006), is removed  
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38 533 from Elegestolepidida for not demonstrating recognisable stages of scale crown growth.  
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40 534 As *Frigorilepis* does not develop neck canals (Fig. 10), the polygonal ultrasculptural  
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43 535 pattern of the crown surface it shares with *Kannathalepis* has been used instead as a  
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45 536 character to support its chondrichthyan affinity (Märss, 2006; Märss et al., 2006). Crown  
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47 537 ornamentation is regarded non-diagnostic at higher taxonomic levels (see above) and at  
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50 538 present no further evidence is available to unite *Frigorilepis* with basal chondrichthyans.  
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52 539 As a consequence, the *Elegestolepis*-type of morphogenesis is the only mechanism of  
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54 540 development recognised in monodontode chondrichthyan scales from the Silurian. The  
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57 541 inclusion of *Ellesmereia* into Elegestolepidida also shows that odontode growth has  
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3 542 persisted as a feature of the integumentary skeleton of chondrichthyans at least until the  
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5 543 Early Devonian (Fig. 10). This last known appearance of an elegestolepid species  
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8 544 coincides with a major diversification of chondrichthyans at the base of the Devonian  
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10 545 (Ginter, 2004; Turner, 2004; Grogan et al., 2012) that sees the emergence of taxa with  
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12 546 body cover of non-growing monodontode scales. Some of these species are known  
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14 547 from body fossils and represent examples of the earliest recorded articulated  
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16 548 chondrichthyan remains (Fig. 10; *Lupopsyrus pygmaeus* (Bernacsek and Dineley, 1977;  
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18 549 Hanke and Davis, 2012) and *Obtusacanthus corroconis* (Hanke and Wilson, 2004)).  
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20 550 *Polymerolepis whitei* (Karatajūtė-Talimaa, 1968, 1998; Hanke et al., 2013), is also  
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22 551 added to the above by being identified on the basis of CT data (Andreev, 2014) to  
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24 552 possess body scales with single odontode crowns that are randomly compartmentalized  
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26 553 into chambered spaces. These scales lack the bony base component of the  
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28 554 elegestolepid squamation, which within the Chondrichthyes has only been documented  
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30 555 in scales with growing crowns (either mon- or poly-odontode). Moreover, *Lupopsyrus*  
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32 556 and *Obtusacanthus*, a pair of genera that have been repeatedly recovered as stem  
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34 557 chondrichthyans in recent hypotheses of early gnathostome phylogeny (Brazeau, 2009;  
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36 558 Davis et al., 2012; Zhu et al., 2013; Giles et al., 2015) are resolved as sister taxa to  
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38 559 Elegestolepidida (Fig. 2) and do not possess scale-neck openings of the pulp canal. A  
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40 560 pattern of vascularization where the pulp opens only towards the lower surface of scales  
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42 561 has a homoplastic distribution inside the stem group, and it is also a feature of the  
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44 562 earliest recorded chondrichthyan polyodontode scales (Sansom et al., 1996; 2001;  
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46 563 Donoghue and Sansom, 2002; Andreev et al., 2015).  
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CONCLUSIONS

The original concept of *Elegestolepis*-type scale morphogenesis (Karatajūtė-Talimaa, 1992) is re-interpreted here to feature a stepwise crown growth and neck canal formation as its diagnostic characteristics. The presence of neck canal openings in *Elegestolepis*-type scales is considered to distinguish them from the growing monodontode scales of the Thelodonti (Märss et al., 2007), whereas the absence of basal bone osteons and hard tissue resorption in these taxa are chondrichthyan apomorphies within crown gnathostomes. This implies that the total-group Chondrichthyes has evolved two distinct morphogenetic processes for generation of single odontode scales, one characteristic for the elegestolepids and the other producing the non-growing *Heterodontus*-type scales (sensu Karatajūtė-Talimaa, 1992), known in detail in euselachians. Consequently, the elegestolepid integumentary skeleton is seen to demonstrate one of the early forms of chondrichthyan scale development that are absent from more derived taxa of the clade. It is further speculated that the contribution of osteogenic tissues to elegestolepid scale units represents a phylogenetically basal state in relation to that of taxa with a solely odontogenically derived squamation.

The shared mode of scale morphogenesis unites *Elegestolepis* (Karatajūtė-Talimaa, 1973) with *Ellesmereia* (Vieth, 1980), *Kannathalepis* (Märss and Gagnier, 2001) and *Deltalepis* gen. nov into the newly erected Order Elegestolepidida, extending the known stratigraphic range of elegestolepid taxa from the Lower Silurian (middle Llandovery) to the Lower Devonian (Lochkovian). Furthermore, a division of the Order into two

588 Families is established upon differences in pulp cavity architecture between  
589 *Kannathalepis* and all the other recognised elegestolepid genera.

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FIGURE CAPTIONS

FIGURE 1. Diagrammatic representation of monodontode scale types in **A**, the Thelodonti and (**B**, **C**) the Chondrichthyes. **A**, a *Thelodus calvus* scale (adapted from Märss and Karatajūtė-Talimaa 2002: fig. 15F) exemplifying the thelodont morphogenetic type; **B**, the *Elegestolepis* morphogenetic type represented by an *Elegestolepis grossi* scale (BU5284); **C**, the *Heterodontus* morphogenetic type represented by a *Triakis semifasciata* scale (BU5341). **Color-coded tissues**: **blue**, enameloid; **brown**, dentine; **gold**, bone. (2/3rd of a whole page width)

FIGURE 2. Distribution of relevant to the study scale characters among select groups of Paleozoic gnathostomes. Tree topology reconstructed from published phylogenies of total-group Chondrichthyes (Grogan et al., 2012) and vertebrates (Sire et al., 2009; Giles et al., 2013, 2015), with the position of Elegestolepidida on the chondrichthyan branch determined from yet to be published analysis by Andreev et al. (representative tree generated in TNT version 1.1 (Goloboff et al., 2008) using a data matrix of 68 equally weighted scale-based characters and 49 Paleozoic jawed-gnathostome taxa). (whole page width)

FIGURE 3. Line drawings depicting the range of crown-surface morphologies in elegestolepid scales. **A**, *Elegestolepis grossi* (BU5284); **B**, *Ellesmereia schultzei*

991 (adapted from Vieth 1980:pl. 9.2); **C**, *Deltalepis magna* (holotype BU5269); **D**,  
 992 *Deltalepis parva* (holotype BU5275). Anterior towards the bottom. (column width)  
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 994 FIGURE 4. Scales of *Elegestolepis grossi* from the Baital Formation of Tuva, Russian  
 995 Federation; ontogenetically mature scales shown in **A**, antero-lateral (BU5285), **B**,  
 996 lateral-crown (BU5285), **C**, lateral (BU5286) and (**D**, BU5286), (**E**, BU5287) crown  
 997 views. **F**, postero-lateral view of BU5289 showing the single neck canal opening of the  
 998 scale crown; **G**, postero-basal view of an ontogenetically young scale (BU5343) with not  
 999 fully formed pedicle support; **H**, basal view of a scale (BU5343) with pedicle support at  
 1000 an advanced stage of formation; **I**, mature scale (BU5289) in basal view exhibiting  
 1001 bulbous basal bone. SEM micrographs. Anterior towards right in (B), towards left in (C),  
 1002 towards the bottom in (D, E) and towards the top in (H, I); arrows indicate neck canal  
 1003 openings, arrowhead indicates the basal opening of the main pulp canal. Scale bars  
 1004 represent 200 µm in (A–E, G, H) and 100 µm in (F, I). (whole page width)

1006 FIGURE 5. Hard tissue structure of *Elegestolepis grossi* scales from the Baital  
 1007 Formation of Tuva, Russian Federation. **A**, vertical cross section of a scale (BU5290) in  
 1008 early stage of bony base formation, etched in 0.5% chromium sulphate solution for 2  
 1009 hours; **B**, detail of **A**, showing the upper medial portion of the crown; **C**, vertical  
 1010 longitudinal section of a scale (BU5291) in advanced stage of basal bone developed  
 1011 (ontogenetically old), etched in 0.5% orthophosphoric acid for 10 minutes; **D**, detail of  
 1012 BU5291 depicting the lower posterior margin of the crown; **E**, detail of the anterior



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3 1013 portion of the crown of BU5291; **F**, vertical transverse section of an ontogenetically old  
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5 1014 scale (BU5292); **G**, basal bone of ontogenetically old scale (BU5293) in vertical  
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8 1015 longitudinal section. (B–E) SEM micrographs; (A, F, G) Nomarski interference contrast  
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10 1016 micrographs. Anterior towards the right in (C–E, G); (B), base; arrowheads in (B–E)  
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12 1017 demarcate the extent of artificially altered dentine, asterisks in (G) denote the borders of  
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14 1018 depositional bone lamellae. Scales bars represent 100  $\mu\text{m}$  in (A, C, F, G) 50  $\mu\text{m}$  in (B,  
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16 1019 E) and 20  $\mu\text{m}$  in (D). (whole page width)  
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24 1021 FIGURE 6. Scales of *Deltalepis magna* gen. et sp. nov. from the Chargat Formation of  
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26 1022 north-western Mongolia. Holotype specimen (BU5269, scale with a five-lobed crown  
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28 1023 and a gracile neck) in **A**, anterior, **B**, antero-lateral and **C**, crown view. **D**, scale  
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30 1024 (BU5270) with gracile neck in basal view. Scales with three-lobe crowns in **E**, anterior,  
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32 1025 **F**, posterior, **G**, lateral (E–G, BU5273) and **H**, crown (BU5271) views. **I**, BU5273 in basal  
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34 1026 view revealing the lower pedicle surface; **J**, basal view of a scale (BU5272) with fully  
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36 1027 formed pedicle support. (A–C, H–J) SEM micrographs; (D–G) volume renderings.  
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38 1028 Anterior towards the right in (B), towards the bottom in (C, H) towards the top in (D, I, J);  
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40 1029 arrow indicates a neck canal opening. Scale bars represent 200  $\mu\text{m}$ . (whole page width)  
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48 1031 FIGURE 7. Scales of *Deltalepis parva* gen. et sp. nov. from the Chargat Formation of  
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50 1032 north-western Mongolia. Holotype (BU5275) in **A**, crown and **B**, anterior-crown view.  
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52 1033 Scale (BU5280) with a gracile neck in **C**, anterior and **D**, posterior view. Scale (BU5277)  
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54 1034 in **E**, anterior and **F**, crown view. **G**, scale (BU5278) with a gracile neck in basal view,  
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1035 exposing the rami of the pulp canal system. Scale (BU5279) with formed pedicle  
1036 support in **H**, basal and **I**, postero-basal view. (A, B, E–I) SEM micrographs; (C, D)  
1037 volume renderings. Anterior towards the bottom in (A, F) towards the top in (G–I);  
1038 arrows indicate neck canal openings, arrowhead indicates the basal opening of the  
1039 main pulp canal. Scale bars represent 200  $\mu\text{m}$  in (A–D, G) and 100  $\mu\text{m}$  in (E, F, H, I).  
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1042 FIGURE 8. Hard tissue structure of *Deltalepis* gen. nov. **A**, longitudinal tomographic  
1043 slice of a *Deltalepis magna* scale (BU5273); **B**, detail of the dentine tissue at the upper  
1044 anterior margin of the crown of a longitudinally sectioned *Deltalepis magna* scale  
1045 (BU5274); **C**, longitudinal tomographic slice of a *Deltalepis parva* scale (BU5280); **D**,  
1046 view of the posterior portion of a *Deltalepis parva* scale (BU5282) crown immersed in  
1047 clove oil. (B, D) Nomarski interference contrast micrographs; (A, C) volume renderings.  
1048 Anterior towards the right in (A) and towards the left in (C). Scale bars represent 100  $\mu\text{m}$   
1049 in (A, C, D) and 50  $\mu\text{m}$  in (B). (whole page width)

1051 FIGURE 9. Volume renderings of the scale canal system (in red) of examined  
1052 elegestolepids. The scales are made translucent in all renderings, with the exception of  
1053 (G). **A–C**, *Elegestolepis grossi* scale (BU5284) from the Baital Formation of Tuva  
1054 (Russian Federation) in **A**, anterior, **B**, postero-lateral and **C**, crown (depicting the lower  
1055 portion of the specimen that is transversely sliced through the neck region) view. **D–F**,  
1056 *Deltalepis magna* scale (BU5273) from the upper Llandovery–lower Wenlock of north

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3 1057 western Mongolia in **D**, crown and **E**, posterior view and a **F**, crown view of the lower  
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6 1058 portion of the same specimen sliced through the neck region. **G–J**, *Deltalepis parva*  
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8 1059 specimens (BU5280 and BU5281) from the upper Llandovery–lower Wenlock of north  
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10 1060 western Mongolia; **G**, BU5280 sliced transversely through the crown in crown view; **H**,  
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12 1061 BU5280 in anterior view; **I**, **J**, BU5281 in **I** posterior and **J**, postero-lateral view. Anterior  
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15 1062 towards the left in (B), towards the top in (C, F, G) and towards the bottom in (D);  
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17 1063 arrows indicate neck canal openings, arrowheads point at the basal opening of the main  
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20 1064 pulp canal. Scale bars represent 100 µm. (whole page width)  
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26 1066 FIGURE 10. Characteristics of monodontode scales of recognised lower Paleozoic  
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28 1067 chondrichthyans and their stratigraphic range. Pink rectangle designates elegestolepid  
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30 1068 taxa. *Elegestolepis* (Karatajūtė-Talimaa, 1973 and data from this study), *Deltalepis*  
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33 1069 (data from this study), *Kannathalepis* (Märss and Gagnier, 2001), *Ellesmereia* (Vieth,  
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35 1070 1980), *Frigorilepis* (Märss et al., 2002, 2006), *Polymerolepis* Karatajūtė-Talimaa, 1998;  
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38 1071 Hanke et al., 2013), *Lupopsyrus* and *Obtusacanthus* (Hanke and Wilson, 2004; Hanke  
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40 1072 and Davis, 2012). (whole page width)  
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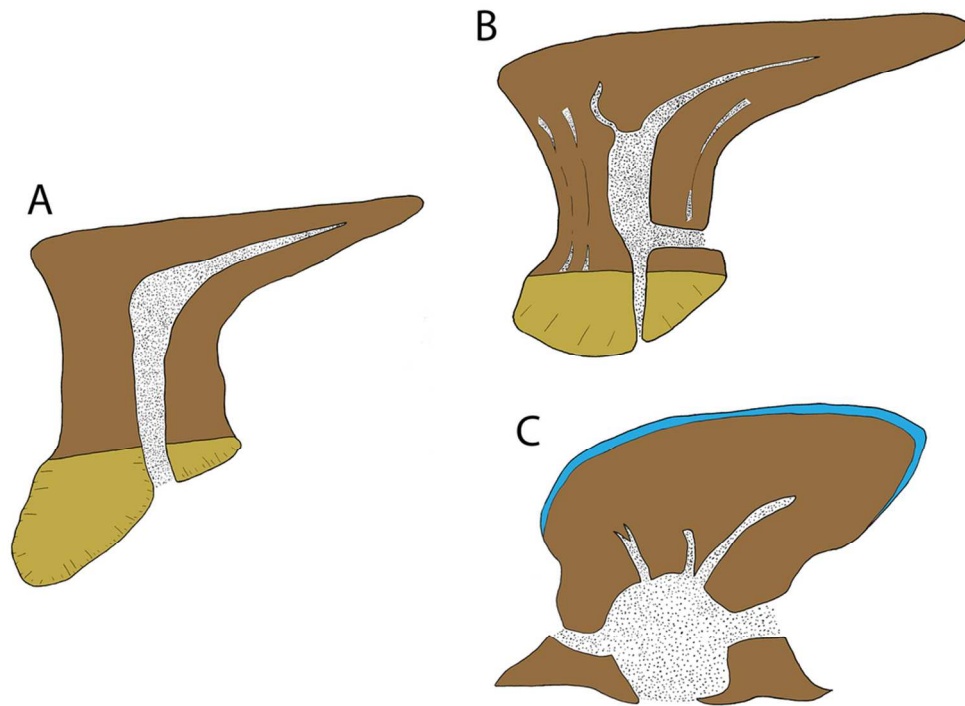


FIGURE 1. Diagrammatic representation of monodontode scale types in A, the Thelodonti and (B, C) the Chondrichthyes. A, a *Thelodus calvus* scale (adapted from Märss and Karatajūtė-Talimaa 2002: fig. 15F) exemplifying the thelodont morphogenetic type; B, the *Egestolepis* morphogenetic type represented by an *Egestolepis grossi* scale (BU5284); C, the *Heterodontus* morphogenetic type represented by a *Triakis semifasciata* scale (BU5341). Color-coded tissues: blue, enameloid; brown, dentine; gold, bone. [2/3 of a whole page width]

Fig1

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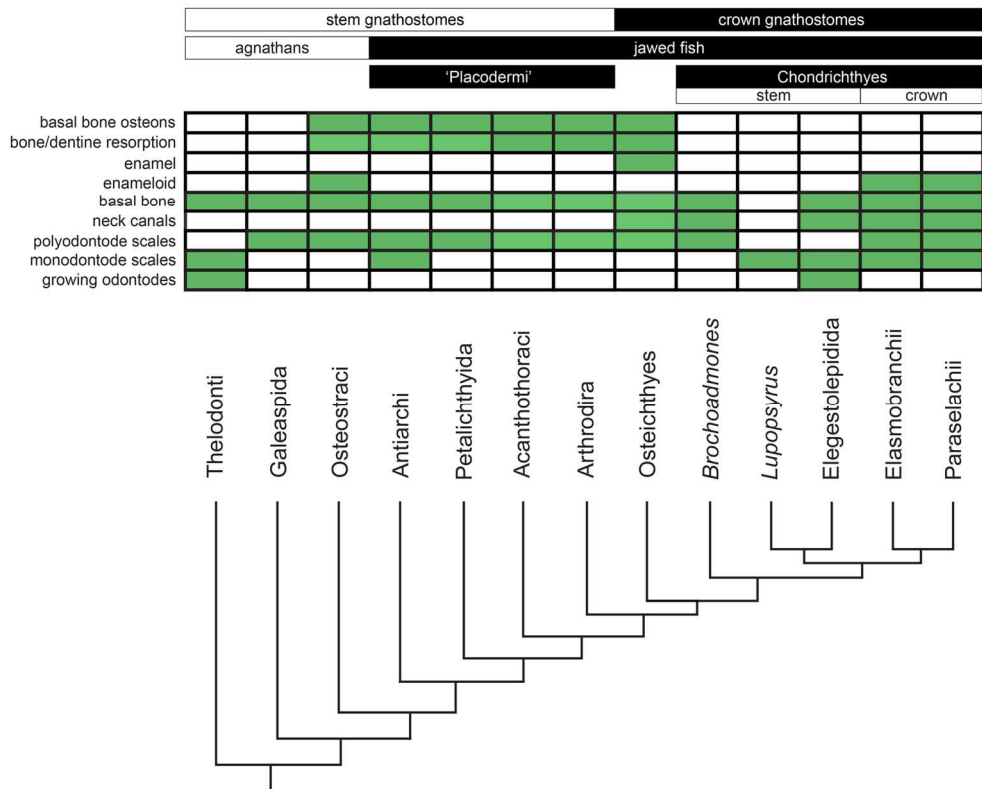


FIGURE 2. Distribution of relevant to the study scale characters among select groups of Paleozoic gnathostomes. Tree topology reconstructed from published phylogenies of total-group Chondrichthyes (Grogan et al., 2012) and vertebrates (Sire et al., 2009; Giles et al., 2013, 2015), with the position of Elegestolepidida on the chondrichthyan branch determined from yet to be published analysis by Andreev et al. (representative tree generated in TNT version 1.1 (Goloboff et al., 2008) using a data matrix of 68 equally weighted scale-based characters and 49 Paleozoic jawed-gnathostome taxa). [whole page width]  
Fig2  
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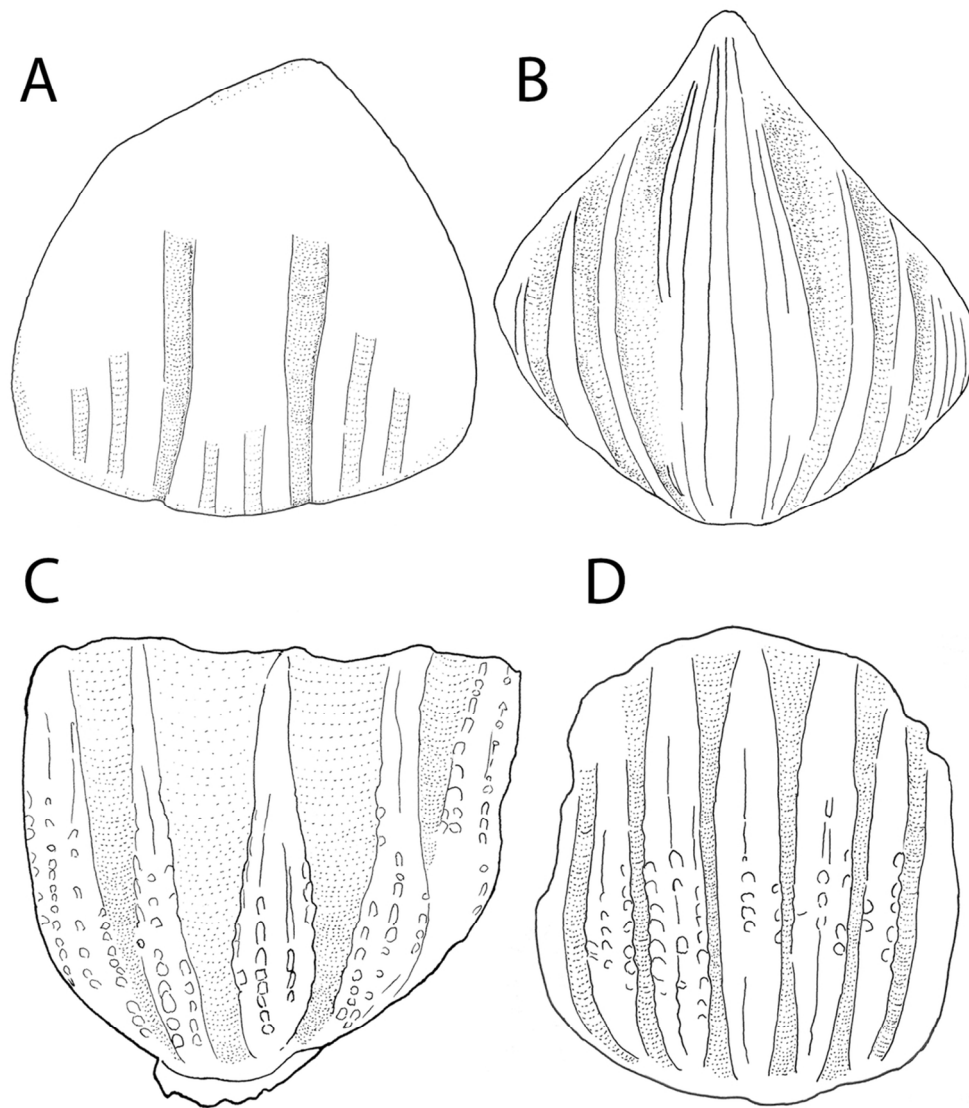


FIGURE 3. Line drawings depicting the range of crown-surface morphologies in elegestolepid scales. A, *Elegestolepis grossi* (BU5284); B, *Ellesmereia schultzei* (adapted from Vieth 1980:pl. 9.2); C, *Deltalepis magna* (holotype BU5269); D, *Deltalepis parva* (holotype BU5275). Anterior towards the bottom. (column width)!! †

Fig. 3

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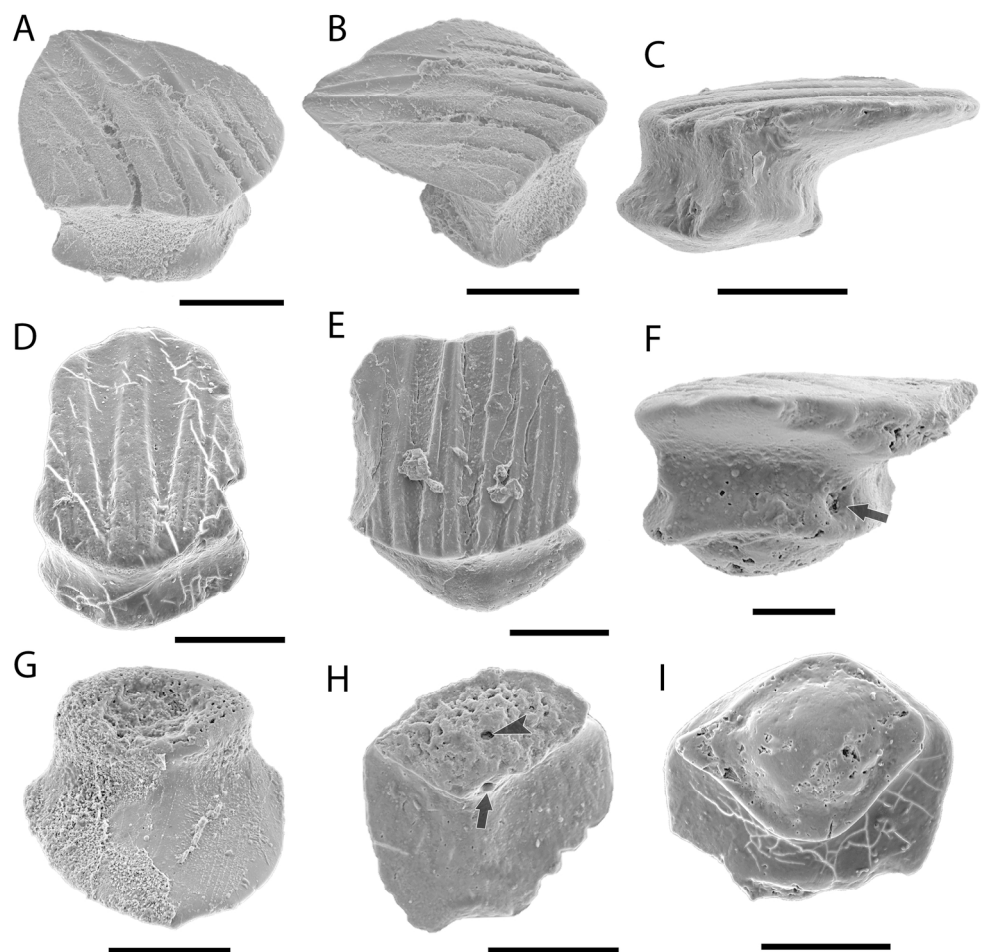


FIGURE 4. Scales of *Elegestolepis grossi* from the Baital Formation of Tuva, Russian Federation; ontogenetically mature scales shown in A, antero-lateral (BU5285), B, lateral-crown (BU5285), C, lateral (BU5286) and (D, BU5286), (E, BU5287) crown views. F, postero-lateral view of BU5289 showing the single neck canal opening of the scale crown; G, postero-basal view of an ontogenetically young scale (BU5343) with not fully formed pedicle support; H, basal view of a scale (BU5343) with pedicle support at an advanced stage of formation; I, mature scale (BU5289) in basal view exhibiting bulbous basal bone. SEM micrographs. Anterior towards right in (B), towards left in (C), towards the bottom in (D, E) and towards the top in (H, I); arrows indicate neck canal openings, arrowhead indicates the basal opening of the main pulp canal. Scale bars represent 200  $\mu$ m in (A–E, G, H) and 100  $\mu$ m in (F, I). [whole page width]

Fig. 4  
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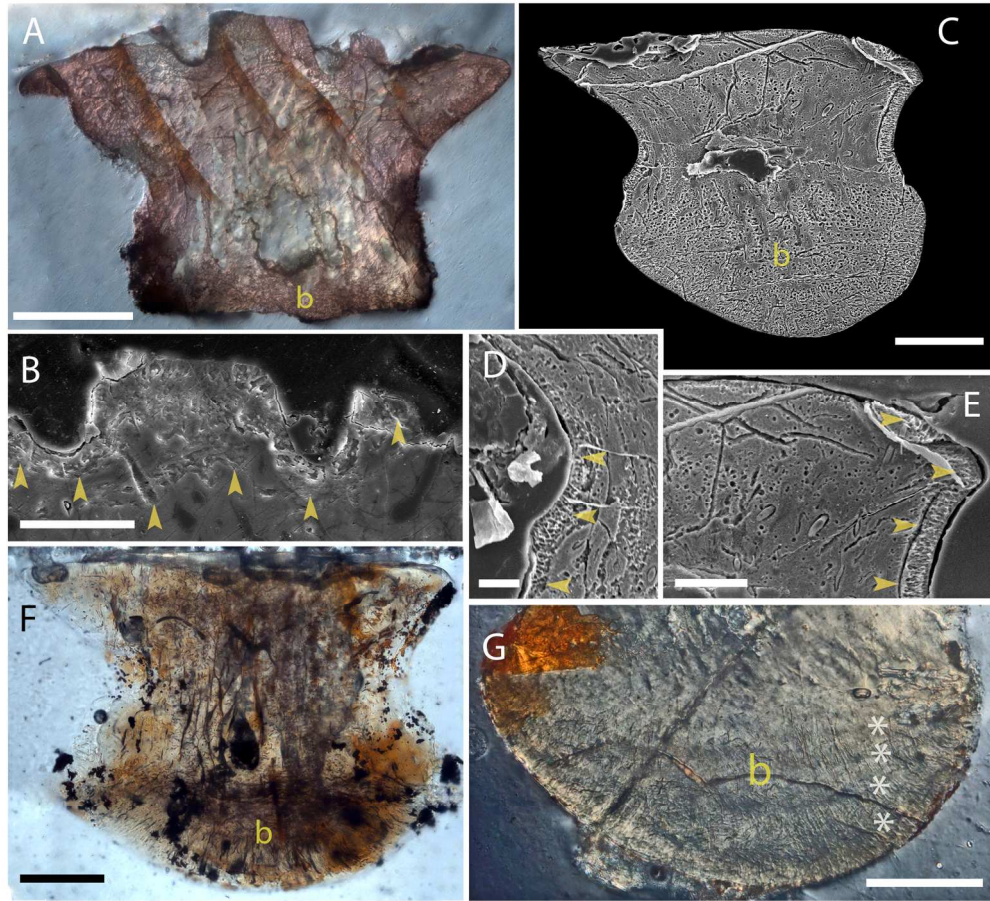


FIGURE 5. Hard tissue structure of *Elegestolepis grossi* scales from the Baital Formation of Tuva, Russian Federation. A, vertical cross section of a scale (BU5290) in early stage of bony base formation, etched in 0.5% chromium sulphate solution for 2 hours; B, detail of A, showing the upper medial portion of the crown; C, vertical longitudinal section of a scale (BU5291) in advanced stage of basal bone developed (ontogenetically old), etched in 0.5% orthophosphoric acid for 10 minutes; D, detail of BU5291 depicting the lower posterior margin of the crown; E, detail of the anterior portion of the crown of BU5291; F, vertical transverse section of an ontogenetically old scale (BU5292); G, basal bone of ontogenetically old scale (BU5293) in vertical longitudinal section. (B–E) SEM micrographs; (A, F, G) Nomarski interference contrast micrographs. Anterior towards the right in (C–E, G); (B), base; arrowheads in (B–E) demarcate the extent of artificially altered dentine, asterisks in (G) denote the borders of depositional bone lamellae. Scale bars represent 100 µm in (A, C, F, G) 50 µm in (B, E) and 20 µm in (D). [whole page width]

Fig. 5  
164x149mm (300 x 300 DPI)



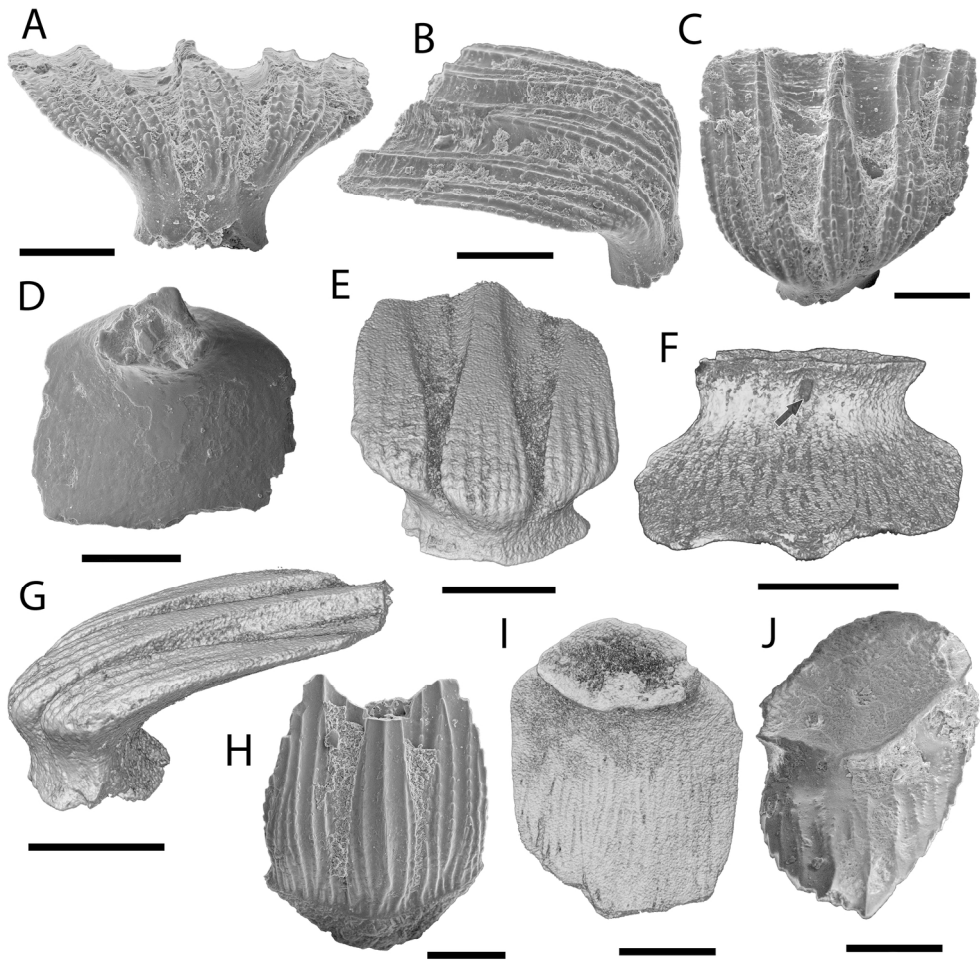


FIGURE 6. Scales of *Deltalepis magna* gen. et sp. nov. from the Chargat Formation of north-western Mongolia. Holotype specimen (BU5269, scale with a five-lobed crown and a gracile neck) in A, anterior, B, antero-lateral and C, crown view. D, scale (BU5270) with gracile neck in basal view. Scales with three-lobe crowns in E, anterior, F, posterior, G, lateral (E–G, BU5273) and H, crown (BU5271) views. I, BU5273 in basal view revealing the lower pedicle surface; J, basal view of a scale (BU5272) with fully formed pedicle support. (A–C, H–J) SEM micrographs; (D–G) volume renderings. Anterior towards the right in (B), towards the bottom in (C, H) towards the top in (D, I, J); arrow indicates a neck canal opening. Scale bars represent 200 μm. (whole page width)

Fig. 6  
177x173mm (300 x 300 DPI)

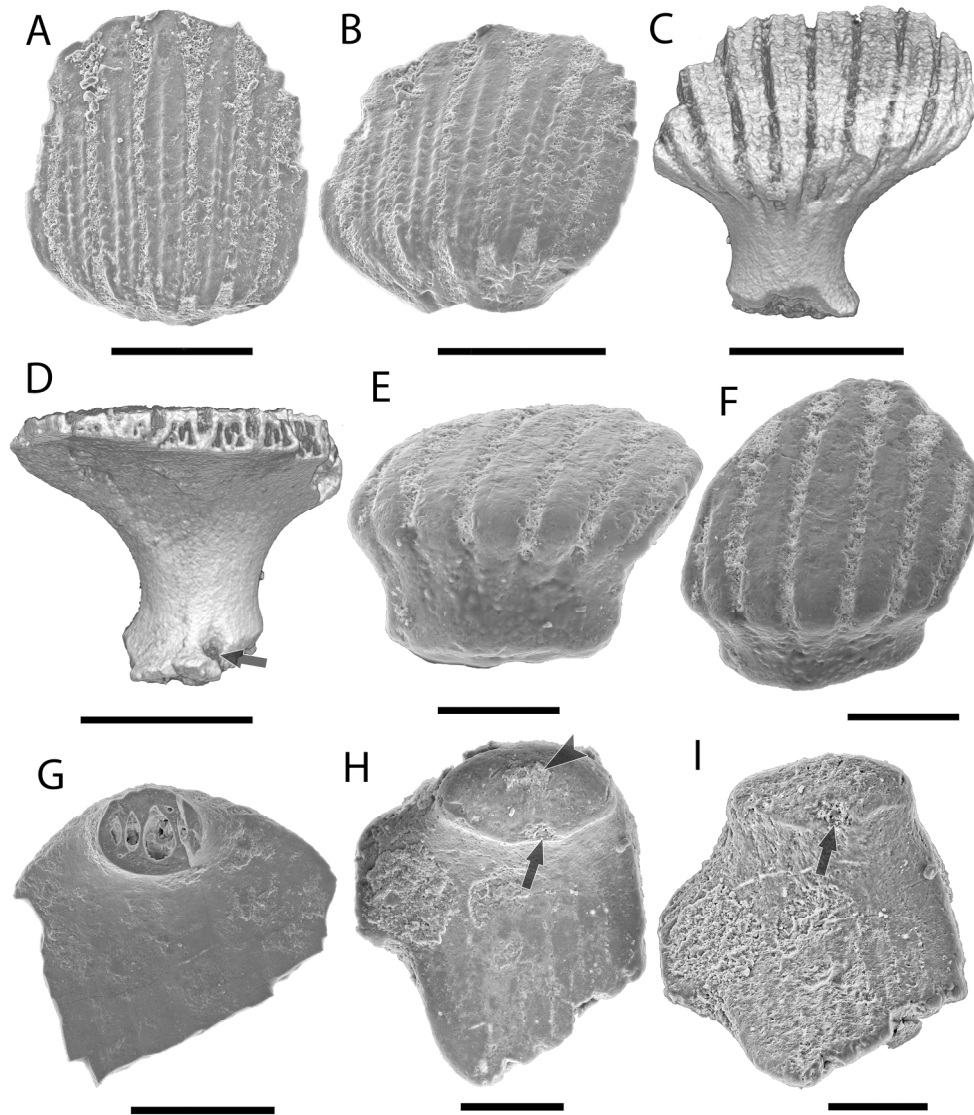


FIGURE 7. Scales of *Deltalepis parva* gen. et sp. nov. from the Chargat Formation of north-western Mongolia. Holotype (BU5275) in A, crown and B, anterior-crown view. Scale (BU5280) with a gracile neck in C, anterior and D, posterior view. Scale (BU5277) in E, anterior and F, crown view. G, scale (BU5278) with a gracile neck in basal view, exposing the rami of the pulp canal system. Scale (BU5279) with formed pedicle support in H, basal and I, postero-basal view. (A, B, E-I) SEM micrographs; (C, D) volume renderings.

Anterior towards the bottom in (A, F) towards the top in (G-I); arrows indicate neck canal openings, arrowhead indicates the basal opening of the main pulp canal. Scale bars represent 200  $\mu$ m in (A-D, G) and 100  $\mu$ m in (E, F, H, I). [whole page width]

Fig. 7  
204x229mm (300 x 300 DPI)

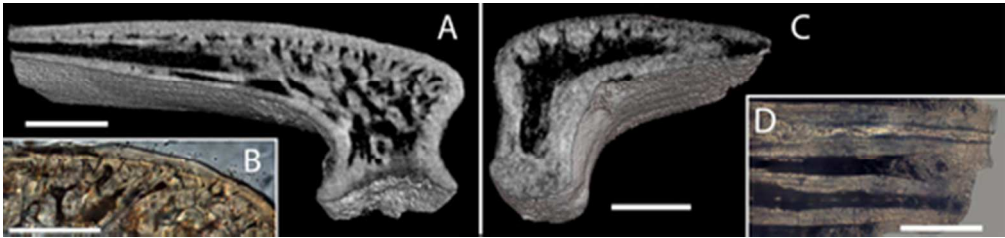


FIGURE 8. Hard tissue structure of *Deltalepis* gen. nov. A, longitudinal tomographic slice of a *Deltalepis magna* scale (BU5273); B, detail of the dentine tissue at the upper anterior margin of the crown of a longitudinally sectioned *Deltalepis magna* scale (BU5274); C, longitudinal tomographic slice of a *Deltalepis parva* scale (BU5280); D, view of the posterior portion of a *Deltalepis parva* scale (BU5282) crown immersed in clove oil. (B, D) Nomarski interference contrast micrographs; (A, C) volume renderings. Anterior towards the right in (A) and towards the left in (C). Scale bars represent 100  $\mu\text{m}$  in (A, C, D) and 50  $\mu\text{m}$  in (B).  
[whole page width]

Fig. 8  
42x9mm (300 x 300 DPI)

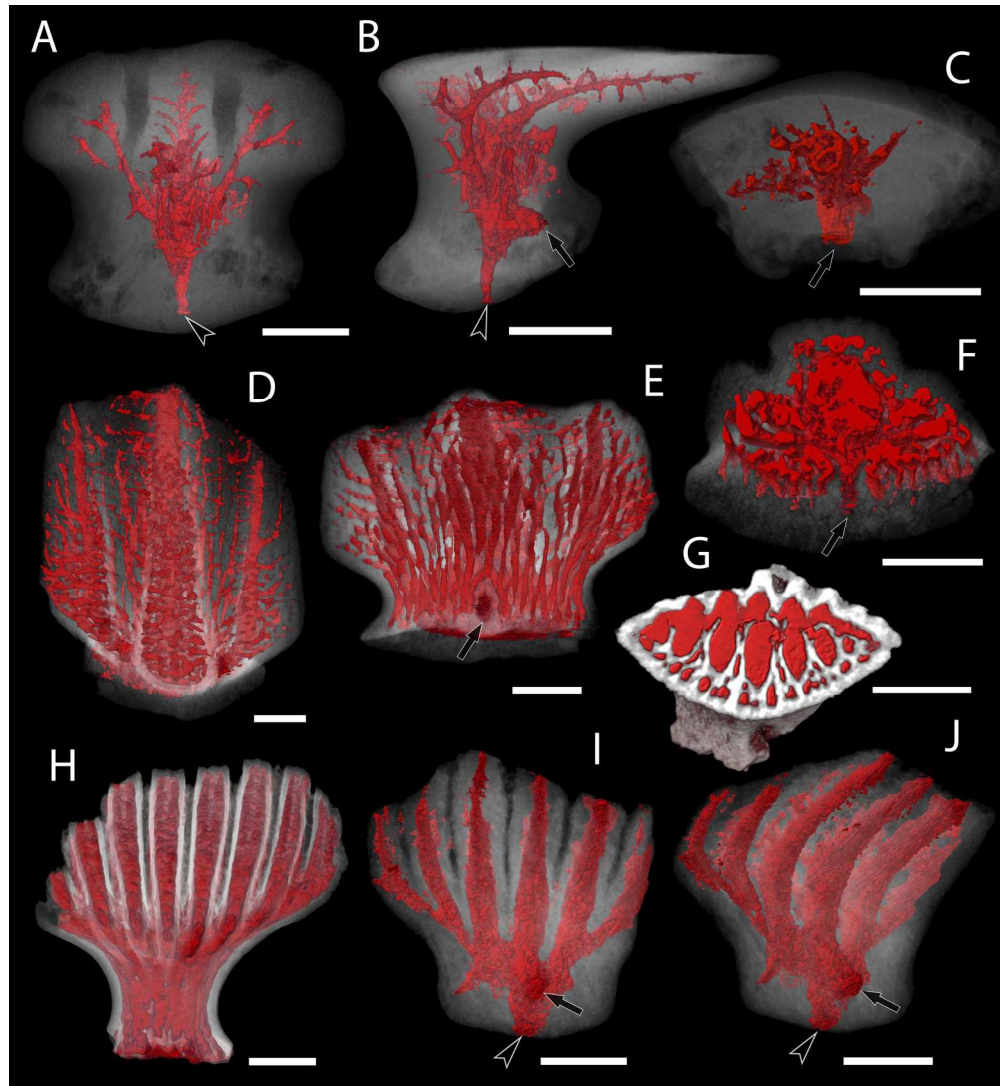


FIGURE 9. Volume renderings of the scale canal system (in red) of examined elegestolepids. The scales are made translucent in all renderings, with the exception of (G). A–C, *Elegestolepis grossi* scale (BU5284) from the Baital Formation of Tuva (Russian Federation) in A, anterior, B, postero-lateral and C, crown (depicting the lower portion of the specimen that is transversely sliced through the neck region) view. D–F, *Deltalepis magnus* scale (BU5273) from the upper Llandovery–lower Wenlock of north western Mongolia in D, crown and E, posterior view and a F, crown view of the lower portion of the same specimen sliced through the neck region. G–J, *Deltalepis parvus* specimens (BU5280 and BU5281) from the upper Llandovery–lower Wenlock of north western Mongolia; G, BU5280 sliced transversely through the crown in crown view; H, BU5280 in anterior view; I, J, BU5281 in I posterior and J, postero-lateral view. Anterior towards the left in (B), towards the top in (C, F, G) and towards the bottom in (D); arrows indicate neck canal openings, arrowheads point at the basal opening of the main pulp canal. Scale bars represent 100 µm. [whole page width]

Fig. 9  
196x212mm (300 x 300 DPI)

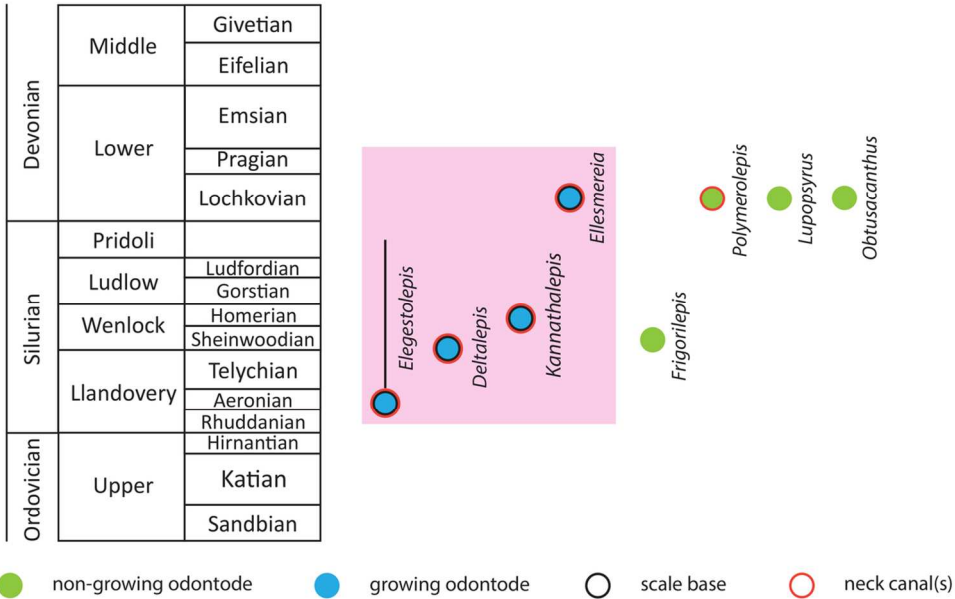


FIGURE 10. Characteristics of monodontode scales of recognised lower Paleozoic chondrichthyans and their stratigraphic range. Pink rectangle designates egestolepid taxa. *Egestolepis* (Karatajūtė-Talimaa, 1973 and data from this study), *Deltalepis* (data from this study), *Kannathalepis* (Märss and Gagnier, 2001), *Ellesmereia* (Vieth, 1980), *Frigorilepis* (Märss et al., 2002, 2006), *Polymerolepis* Karatajūtė-Talimaa, 1998; Hanke et al., 2013), *Lupopsyrus* and *Obtusacanthus* (Hanke and Wilson, 2004; Hanke and Davis, 2012).  
[whole page width]

Fig. 10  
115x72mm (300 x 300 DPI)